



12-10-01 ~~11/10~~ 0360

INFORMATION DISCLOSURE CITATION (Use several sheets if necessary)	ATTY. DOCKET NO.	SERIAL NO.
	96606/16UTL	09/765,061 /
	APPLICANT Melanie M. Sohocki, et al.	
	FILING DATE 1/17/2001	GROUP

#8
Supp
6-30-02

U.S. PATENT DOCUMENTS

EXAMINER	DOCUMENT NO.	DATE	NAME	CLASS	SUBCLASS	FILING DATE
INITIAL						IF APPROPRIATE

FOREIGN PATENT DOCUMENTS

DOCUMENT NO.	DATE	COUNTRY	CLASS	SUBCLASS	TRANSLATION
					YES NO

OTHER DOCUMENTS (Including Author, Title, Date, Pertinent Papers, Etc.)

	3/2000	Investigative Ophthalmology & Visual Science- Article entitled "A Novel Locus for Leber Congenital Amaurosis (LCA4) with Anterior Keratoconus Mapping to Chromosome 17p13" by Abdul Hameed et al.
	1/2000	Nature Genetics- Article entitled "Mutations in a New Photoreceptor-Pineal Gene on 17p Cause Leber Congenital Amaurosis" by Melanie M. Sohocki et al.
	8/2000	Human Mutation 17:42-51- Article entitled "Prevalence of Mutations Causing Retinitis Pigmentosa and Other Inherited Retinopathies" by Melanie M. Sohocki et al.
	3/2/1999	Genomics 58, 29-33- Article entitled "Localization of Retina/Pineal-Expressed Sequences: Identification of Novel Candidate Genes for Inherited Retinal Disorders" by Melanie M. Sohocki et al.
	5/3/2000	Molecular Genetics and Metabolism 70, 142-150- Article entitled "Prevalence of AIPL1 Mutations in Inherited Retinal Degenerative Disease" by Melanie M. Sohocki et al.
	3/5/2001	Mammalian Genome 12, 566-568- Article entitled "Comparative Analysis of Aryl-Hydrocarbon Receptor Interacting Protein-Like 1 (Aipl1), a Gene Associated With Inherited Retinal Disease in Humans" by Melanie M. Sohocki et al.
	1999	American Journal of Human Genetics, 65:A112, 1999- Article entitled "Human Aryl-Hydrocarbon Interacting Protein-Like 1 Gene (AIPL1), a Candidate for Inherited Retinal Disorders: Mapping to 17p13, Characterization and Mutation Testing" by Melanie M. Sohocki et al.
	2000	Investigative Ophthalmology & Visual Science, 41:S94,2000- Article entitled "Mutations in AIPL1, A Novel Photoreceptor/Pineal-Expressed Gene on 17p13, Cause Leber Congenital Amaurosis (LCA4)" by Melanie M. Sohocki et al.
	2001	Investigative Ophthalmology & Visual Science, 42:S645,2001- Article entitled "Role of AIPL1 in LCA and Related Inherited Retinal Diseases", by Melanie M. Sohocki et al.
	2000	American Journal of Human Genetics, 67:388S,2000-Article entitled "Comparative Sequencing of Aryl-Hydrocarbon Interacting Protein Like-1 (AIPL1), A Protein Associated with Leber Congenital Amaurosis" by Melanie M. Sohocki et al.
	2000	American Journal of Human Genetics, 67:411S,2000- Article entitled "Molecular Studies of AIPL1, a Gene Causing Leber Congenital Amaurosis" by D.L. Tirpak et al.
	2001	Investigative Ophthalmology & Visual Science, 42:S655,2001- Article entitled "Yeast Two-Hybrid Analysis of AIPL1-Binding Proteins" by D.L. Tirpak et al.
EXAMINER		DATE CONSIDERED

CERTIFICATE OF MAILING BY "EXPRESS MAIL" (37 CFR 1.10)Applicant(s): **Melanie M. Sohocki, et al.**

Docket No.

96606/16UTL

Serial No.

09/765,061

Filing Date

01/17/2001

Examiner

Group Art Unit

Invention: **Mutations in a Novel Photoreceptor-Pineal Gene on 17P Cause Leber Congenital Amaurosis (LCA 4)**I hereby certify that this **Information Disclosure Statement***(Identify type of correspondence)*

is being deposited with the United States Postal Service "Express Mail Post Office to Addressee" service under 37 CFR 1.10 in an envelope addressed to: The Assistant Commissioner for Patents, Washington, D.C. 20231

on **November 5, 2001***(Date)***Sandra Cortinas***(Typed or Printed Name of Person Mailing Correspondence)*

A handwritten signature in cursive script that reads "Sandra Cortinas".

*(Signature of Person Mailing Correspondence)***EV 008 608 327 US***("Express Mail" Mailing Label Number)***Note: Each paper must have its own certificate of mailing.**